L Number	Hits	Search Text		
1	2711	interphase	DB	Time stamp
1	2/11	Incerphase	USPAT;	2002/08/09 06:26
2	16904	chromosome	US-PGPUB	
-	10004	CITOMOSOME	USPAT;	2002/08/09 06:26
4	1100839	damag\$8 or break\$8 or fragment\$8	US-PGPUB	
•	1100035	damagso of breakso of fragmentss	USPAT;	2002/08/09 06:27
5	584840	mitogen or bleomycin or mms or	US-PGPUB	
	304040	methanesulfonate or arac	USPAT;	2002/08/09 06:29
6	452		US-PGPUB	
	452	break\$8 or fragment\$8) and (mitogen or	USPAT;	2002/08/09 06:29
		bleomycin or mms or methanesulfonate or arac	US-PGPUB	
		)		
7	4	interphase same chromosome same (damag\$8 or	USPAT;	2002/00/00 06 00
		break\$8 or fragment\$8) same (mitogen or	US-PGPUB	2002/08/09 06:29
		bleomycin or mms or methanesulfonate or arac	US-FGFUB	
		)		
8	22750	mitogen or bleomycin or methanesulfonate or	USPAT;	2002/08/09 06:29
		arac	US-PGPUB	2002/08/09 08:29
9	87	interphase and chromosome and (damag\$8 or	USPAT;	2002/08/09 06:30
		break\$8 or fragment\$8) and (mitogen or	US-PGPUB	2002/00/03 00:30
		bleomycin or methanesulfonate or arac )	10102	
10	49	interphase and (chromosome same (damag\$8 or	USPAT;	2002/08/09 06:30
}		break\$8 or fragment\$8)) and (mitogen or	US-PGPUB	,,
		bleomycin or methanesulfonate or arac )		
11	26		USPAT;	2002/08/09 06:34
		fragment\$8)) same (mitogen or bleomycin or	US-PGPUB	
		methanesulfonate or arac )		
12	684	The same same taxing to or prouted or	USPAT;	2002/08/09 06:35
		fragment\$8)) and (mitogen or bleomycin or	US-PGPUB	
		methanesulfonate or arac ) and (alzheimer\$3		
13	5550	or cancer\$9)		
13	5759	chromosome same (damag\$8 or break\$8 or	USPAT;	2002/08/09 06:35
14	841	fragment\$8)	US-PGPUB	
	041	(chromosome same (damag\$8 or break\$8 or	USPAT;	2002/08/09 06:35
		fragment\$8)) and (mitogen or bleomycin or methanesulfonate or arac )	US-PGPUB	
15	684	((chromosome same (damag\$8 or break\$8 or		
13	004	fragment\$8)) and (mitogen or bleomycin or	USPAT;	2002/08/09 06:35
		methanesulfonate or arac )) and (alzheimer\$	US-PGPUB	:
		or cancer\$)		İ
16	255		USPAT;	2002/00/00 05 25
		fragment\$8)) and (mitogen or bleomycin or	US-PGPUB	2002/08/09 06:35
		methanesulfonate or arac )) and (alzheimer\$)	OD-EGFOR	
17	180	(((chromosome same (damag\$8 or break\$8 or	USPAT;	2002/08/09 06:36
		fragment\$8)) and (mitogen or bleomycin or	US-PGPUB	2002,00,00 00.30
		methanesulfonate or arac )) and	00 10102	
		(alzheimer\$)) and metaphas\$		
19	317		USPAT;	2002/08/09 06:59
		fragment\$8)) and (mitogen or bleomycin or	US-PGPUB	, , ,
		methanesulfonate or arac )) and (alzheimer\$		
		or cancer\$)) and (dNTP or datp or dttp or		
21	2.0	dctp or dgtp)	-	
21	20	(((chromosome same (damag\$8 or break\$8 or	USPAT;	2002/08/09 07:15
		fragment\$8)) and (mitogen or bleomycin or	US-PGPUB	
		methanesulfonate or arac )) and (alzheimer\$		
22	1453	or cancer\$)) and deoxynucleotidyl		
	1403	deoxynucleotidyl or apotag or tunel	USPAT;	2002/08/09 07:15
23	1	(deoxymuoleotidy) or anotae as turned	US-PGPUB	
		(deoxynucleotidyl or apotag or tunel) same (damag\$8 or break\$8 or fragment\$8) same	USPAT;	2002/08/09 07:16
		chromosome	US-PGPUB	
24	144	(deoxynucleotidyl or apotag or tunel) and	IICDAM -	2002/02/22 27 7
		((damag\$8 or break\$8 or fragment\$8) same	USPAT;	2002/08/09 07:17
		chromosome) and (mitogen or bleomycin or mms	US-PGPUB	
		or methanesulfonate or arac )		
25	57	(deoxynucleotidyl or apotag or tunel) and	USPAT;	2002/08/09 07:18
		((damag\$8 or break\$8 or fragment\$8) near5	US-PGPUB	2002/00/09 07:18
		chromosome) and (mitogen or bleomycin or mms		
		or methanesulfonate or arac )	1	

26	9	(doorgranglootidal on contract the last		
20	,	(deoxynucleotidyl or apotag or tunel) and	USPAT;	2002/08/09 07:20
		((damag\$8 or break\$8 or fragment\$8) near5	US-PGPUB	
		chromosome) and (mitogen or bleomycin or		
		methanesulfonate or arac )		
27	32932	(chromosome or DNA) near8 ((damag\$8 or	USPAT;	2002/08/09 07:21
		break\$8 or fragment\$8) or cut)	US-PGPUB	2002/00/03 07:21
28	397		USPAT;	2002/08/09 07:21
		break\$8 or fragment\$8) or cut)) same	US-PGPUB	2002/08/09 07:21
		(deoxynucleotidyl or apotag or tunel)	U3-FGFUB	İ
29	86		TIODAM	2000/00/00
		break\$8 or fragment\$8) or cut)) same	USPAT;	2002/08/09 07:21
		/doorners and and dell are a set	US-PGPUB	
		(deoxynucleotidyl or apotag or tunel)) and		
		(mitogen or bleomycin or methanesulfonate or		
2.0		arac)		
30	18	(((chromosome or DNA) near8 ((damag\$8 or	USPAT;	2002/08/09 07:21
		break\$8 or fragment\$8) or cut)) same	US-PGPUB	
		(deoxynucleotidyl or apotag or tunel)) and		
		interphase		

(FILE 'HOME' ENTERED AT 11:31:03 ON 09 AUG 2002)

FILE 'MEDLINE, BIOSIS, CAPLUS, EMBASE, SCISEARCH' ENTERED AT 11:31:14 ON 09 AUG 2002

L1 31964 S TUNEL OR APOTAG OR TERMINAL (4A) TRANSFERASE OR DEOXYNUCLEOTI

L2 60438 S INTERPHASE L3 134 S L2 AND L1

L4 94 DUP REM L3 (40 DUPLICATES REMOVED)

L5 57 S L1 (P) L2

L6 18 DUP REM L5 (39 DUPLICATES REMOVED)

L7 36847 S L1 OR TDT L8 57 S L7 (P) L2

L9 18 DUP REM L6 (0 DUPLICATES REMOVED)

FILE 'STNGUIDE' ENTERED AT 11:37:51 ON 09 AUG 2002

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L9 ANSWER 15 OF 18 MEDLINE

ACCESSION NUMBER: 93011695 MEDLINE

DOCUMENT NUMBER: 93011695 PubMed ID: 1397093

TITLE: Intracellular localization of terminal transferase during

the cell cycle.

AUTHOR: Di Primio R; Trubiani O; Bollum F J

CORPORATE SOURCE: Istituto di Morfologia Umana Normale, Facolta di Medicina,

Universita di Chieti, Italy.

CONTRACT NUMBER: CA-23262 (NCI)

SOURCE: EXPERIMENTAL CELL RESEARCH, (1992 Oct) 202 (2) 405-11.

Journal code: 0373226. ISSN: 0014-4827.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199210

ENTRY DATE: Entered STN: 19930122

Last Updated on STN: 19930122 Entered Medline: 19921030

AB Changes in the localization of terminal transferase

during the cell cycle in random cultures of human pre-T leukemia line RPMI-8402 were examined by light and electron microscopy on immunoperoxidase-stained preparations. Paraformaldehyde-fixed and saponin-permeabilized human cells were used with a monoclonal anti-human terminal deoxynucleotidyl transferase (TdT)

primary reagent to demonstrate changes in enzyme distribution occurring between **interphase** and mitosis. Nuclear localization is found uniformly during **interphase**. At metaphase, however, the majority

of TdT staining appears randomly distributed in the cytoplasm and traces of TdT staining remain associated with mitotic chromatin. At later phases,

when the daughter cells are forming, the enzyme again appears to be restricted to the new nuclear structure.

L9 ANSWER 10 OF 18 MEDLINE

ACCESSION NUMBER: 96428453 MEDLINE

DOCUMENT NUMBER: 96428453 PubMed ID: 8831556

TITLE: DNA segments sensitive to single-strand-specific nucleases

are present in chromatin of mitotic cells.

AUTHOR: Juan G; Pan W; Darzynkiewicz Z

CORPORATE SOURCE: Cancer Research Institute, New York Medical College,

Valhalla 10595, USA.

CONTRACT NUMBER: RO 28704

SOURCE: EXPERIMENTAL CELL RESEARCH, (1996 Sep 15) 227 (2) 197-202.

Journal code: 0373226. ISSN: 0014-4827.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199611

ENTRY DATE: Entered STN: 19961219

Last Updated on STN: 19961219 Entered Medline: 19961105

AB It was observed before that DNA in situ in chromatin of mitotic cells is more sensitive to denaturation than DNA in chromatin of interphase cells. DNA sensitivity to denaturation, in these studies, was analyzed by exposing cells to heat or acid and using acridine orange (AO), the metachromatic fluorochrome which can differentially stain double-stranded (ds) vs single-stranded (ss) nucleic acids, as a marker of the degree of

DNA denaturation. However, without prior cell treatment with heat or acid no presence of single-stranded DNA in either mitotic or interphase cells was detected by this assay. In the present experiments we demonstrate that DNA in situ in mitotic cells, without any prior treatment

that can induce DNA denaturation, is sensitive to ss-specific S1 and mung bean nucleases. Incubation of permeabilized human T cell leukemic MOLT-4, promyelocytic HL-60, histiomonocytic lymphoma U937 cells, or normal PHA-stimulated lymphocytes with S1 or mung bean nucleases generated extensive DNA breakage in mitotic cells. DNA strand breaks were detected using fluorochrome-labeled triphosphonucleotides in the reaction catalyzed

by exogenous terminal deoxynucleotidyl

transferase. Under identical conditions of the cells' exposure to ss-specific nucleases, DNA breakage in interphase cells was of an order of magnitude less extensive compared to mitotic cells. The data indicate that segments of DNA in mitotic chromosomes, in contrast to interphase cells, may be in a conformation which is sensitive to ss nucleases. This may be a reflection of the differences in the torsional

stress of DNA loops between **interphase** and mitotic chromatin. Namely, greater stress in mitotic loops may lead to formation of the hairpin-loop structures by inverted repeats; such structures are sensitive

to ss nucleases. The present method of detection of such segments appears to be more sensitive than the use of AO. The identification of mitotic cells based on sensitivity of their DNA to ss nucleases provides an additional method for their quantification by flow cytometry.

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